# **Copyright Warning & Restrictions**

The copyright law of the United States (Title 17, United States Code) governs the making of photocopies or other reproductions of copyrighted material.

Under certain conditions specified in the law, libraries and archives are authorized to furnish a photocopy or other reproduction. One of these specified conditions is that the photocopy or reproduction is not to be "used for any purpose other than private study, scholarship, or research." If a, user makes a request for, or later uses, a photocopy or reproduction for purposes in excess of "fair use" that user may be liable for copyright infringement,

This institution reserves the right to refuse to accept a copying order if, in its judgment, fulfillment of the order would involve violation of copyright law.

Please Note: The author retains the copyright while the New Jersey Institute of Technology reserves the right to distribute this thesis or dissertation

Printing note: If you do not wish to print this page, then select "Pages from: first page # to: last page #" on the print dialog screen



The Van Houten library has removed some of the personal information and all signatures from the approval page and biographical sketches of theses and dissertations in order to protect the identity of NJIT graduates and faculty.

# PREPARATION AND HEDICTION OF AZO DIES FROM 2-AMINO PREMOL-4-SULFORIC ACID

BY

OUY J. WILETR

### Submitted in Partial Fulfillment

of the Requirements

for the Degree of

MASTER OF SCIENCE

in Chemical Engineering

in the

Graduate Division

at the

Newark College of Engineering

# ACKINE LE LOSE SENT

The author wished to express his gratitude to Professor

James A. Bradley, of the Newark College of Engineering, for his
counsel and guidance throughout the investigation.

# DART OF CUTTION

									Page
ADSTRACT	*	•	•	•	•	*	•	•	1
to the second second second	MOI	uM.	•	•	•	*	•	•	2
The second second second	•	•	•	•	•	*	•	•	Ç
APPLIANCE	•	•	•	•	•	•	•	•	5
AND VERSELL LINE	•	•	•	•	*	•	•	•	6
	Z:C	T.V.	•	*	•	•	*	•	7
erogore 2		of 2-p nic ac		-aso-yi	henol-	*	•	•	7
Propara di		of 4. 1-6-su					•	¥	9
Propara Th		of 4-n 4-sulf			-(1 asc	) 6) <del></del>	•	•	11
Pedu <b>oti</b> ph	on of enol-	4-nit 4-sulf	ro phe onic a	enyl-(	l azo e		•	•	12
Prepara ph		of nap			azo 6)-		•	•	10
Moduo <b>ti</b> My		naght 4-sulf			o 6) <del>-</del>	•	•	•	14
Propere de		of mag 4-sulf			azo 5)-	•	•		<b>1</b> 5
odneti Ni		nepht aulf			0 6)-	•	•	•	16
Diesoti	sctio	r of 1	-arino	<b>)-2-</b> 20)	phthol	•	•	•	17
Constitution	r 09	1_0:4r	A-9-016	an Interface	1 24000	~~~1 <i>2</i> ~			30

# PARIS OF CHIEFARS

									20 <b>00</b>
Dieze	naphthol	. ond Laisa	coupli Uforic	to gri	l-cuino		#	•	19
Coupl	ing of ?	-amir	o pher	201	sulfonic	a <b>ci</b> d	•	•	£0
Prepa	ration of				0 6)-	•	•	•	21
Poduc	phonol-	raghi Luc-	hyl-(2 onic a	ecić	3)-	•	•	•	20
Prone	ration ( Phonol—				6)-	•	•	•	Ð'#
Dochio	tion of phenol-4	naght L-sulf	hyl-(i	l aso ( acid	5)-	•	•	•	25
	LLRUU	.TC	•	•	•	•	•	•	೭೦
DICOUSTICE	•	•	•	•	•	•	•	•	30
Diago	tication	ı		*	•	•	•	•	<b>3</b> 0
Coupl	ing	*	٠	*	٠	•	•	•	5 <b>1</b>
Reduc	tion	•	•	ð	*	•	•	•	32
STILLIN	•	•	•	•	•		•	•	<b>5</b> 5
FICHBARIAN	Gree			4	_		_		55

### AIR THROOT

Reduction of azo dyes is presented as a nothed for the synthesis of 2-knine phenol-4-sulfenic acid.

The synthesis and reduction of some ase derivatives of 2-amino phonol-4-sulfonic acid are described.

Preparation of the diazonium compounds of aniline. nitro aniline, 1-amino-2-naphthol, 1-amino-2-naphthol-1sulfonic acid, 2-amino phenol-4-sulfonic acid, 1-naphthylamino, and 2-maphthylamine is discussed and their ability
to couple with phenol-4-sulfonic acid evaluated.

Reduction of the resultant azo dyes is discussed with reference to fission and the benzidine rearrangement.

An over-all presentation of results is given in Tables
I. II. and III on pages 27 - 29.

# STATE SENT OF PROBLEM

The main purpose of this study was the development of a synthesis for 2-amino phenol-4-sulfonic acid, using dissonius compounds and phenol-4-sulfonic acid.

#### INTERNATION

2 9

2-Amino phenol-1-sulforic acid is one of the important intermediates used in the manufacture of the mone ase and dihydroxy dycstuffe. From it such dyes (27) as Acid Alizarin Brown B. Falatine Chrome Brown U. Acid Alizarin Garnet B. Acid Alizarin Violet II. and Diamond Black PV are produced.

The industrial process (27) for production of 2-arino phenol-4-sulfonic acid stems from **chlorobenzene** as fellows:

A mothod using the sulfonation of 2-amino-4-phenol is also reported (2).

In addition to reduction of nitro compounds, emines are also prepared by the reduction of also compounds. A search of the literature indicated that this latter procedure has not been successfully applied to the production of 2-nmino phenol-4-sulfonic soid. This

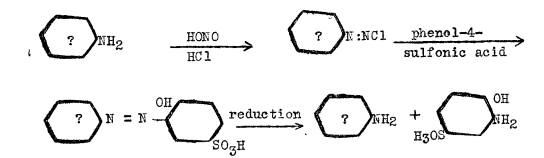
lack sight be secribed to problems associated with:

- 1. Forastion of the diagonium selt.
- 2. Counding.
- 3. Reduction of the eso link to give fission without benzidine rearrangement.
- 4. Isolation of amines from the reduction mixture.

Conditions for dissotisation and coupling vary with the compound and no ettemet will be made to predict procedure. An excellent treatise on the aromatic disso-compounds, their preparation and reactions, has been prepared by Saunders (21).

Reduction of Spentic ase compounds has been the subject of such research. Asines, hydrane compounds, beneidines, or semidines can be formed, depending upon the reduction procedure and the compound to be reduced. Robinson (19) indicates that except for o-emino-ase-bensene, aromatic ase compounds with the para position free reduce to hydrane compounds and rearrange to beneidines in strong acid. Jacobson (16) found that methyl, brene, inde, acetoxy, or ethoxy groups in the para position do not hinder hydrane formation and scidification gives semidines (diphenyl amines) instead of beneidines. In 1937 Shitmore and Revukes (28) presented a new method for reduction of aromatic and compounds -- raney nickel catalyst and hydrogen under pressure -- and were able to reduce and benzene to aniline with a 96% yield.

Forsation and reduction of the aso compound, from which 2-amino phenol-4-sulfonic acid would be produced, might be illustrated thus:



It is evident that the main problems involved are the selection of "amine-?" and the reduction of its aso dye. This investigation deals with the dissenius compounds of aniline, p-nitro aniline, 1-amino-2-naphthol, 1-amino-2-naphthol-4-sulfonic soid, 2-amino phenol-4-sulfonic soid, 1-naphthylamine, and 2-naphthylamine as possibilities.

Phenol-4-sulfonic acid was selected as the acceptor in the coupling reaction with the diasonium selt because:

- Two of the three required functional groups are present and in position.
- 2. Coupling is limited to the ortho position. There are no isomore.
- 3. Pission, by reduction, will yield the desired saine.

### APPARATIE

Except for the Surgess-Parr apparatus (12) used for reductions with raney mickel catalyst, all backers, Clasks, and glas were were of the type that can be purchased in any laboratory supply house.

#### FIRACCIMES

Grade of reagent is indicated in the procedure for each compound. Distilled water was used whenever the use of alkali was required.

Since phenol-4-sulfonic acid was used as the acceptor in all but two of the coupling reactions, it sight be considered a reagent. To evoid repetition, its preparation from sulfanilic acid by a acdified Remarkh (22) procedure is given here, and this preparation applies wherever phenol-4-sulfonic acid was used:

Sulfenilic seid sonohydrate, analytical grade, 9.1 gm. (0.0477 mole), was dissolved in a solution of 2.6 gm. anhydrous sodium carbonate (0.024 sole) in 49 cc. of distilled water, by heating.

U.S.P. sodium mitrite, 3.5 gm., was dissolved in 9.5 cc. distilled water. A.C.S. grade hydrochloric acid, 10.3 cc., was sixed into 57 gm. of ice. The sodium sulfanilate solution was chilled to 15°C., mixed with the sodium nitrite solution, and the whole poured slowly into the rapidly stirred, iced acid. The white, sulfanilic acid diamonium selt precipitated almost at the start of addition. Care was taken to control addition so that hydrolysis did not take place, as indicated by a red color appearing at the mixing point of the two solutions. A positive test for nitrous soid was obtained on the solution with potassium icedide starch paper at the end of addition.

(Sulfanilic acid diamonium salt gave a bright crange color when spot tested with alcoholic, alkaline 2-naphthol).

motality = about 0 C477 x 1002.

- about 0.40

H20

49

117.5

with vigorous stirring, the solution was quickly heated to boiling on an electric hot plate. The total heating time was 8 to 10 minutes. As the temperature rose the dissonium salt dissolved, the solution turned straw yellow, then orange, formed at 65 - 70°C., and continued to fosm until hydrolysis was complete after heating at 100°C. for 2 to 3 minutes. The wine-red solution of phenol-4-sulfonic sold was spot tested for absence of dissonium salt with alkaline 2-maphthol, as noted above, and cooled to 15°C. When used for coupling reactions, this solution was presumed to contain 0.0477 mole of phenol-4-sulfonic acid.

## EXPERIMENTAL PROCEDURE

#### PHERADALICA of 2-PHERMI-ALO-PHERMI-A-SULFONIC ACID

C.P. Amiline, 4.5 gm. (0.048 mole), was diametrized by the method of Schwalbe (23) for p-nitro entities. After mixing, the solution gave a positive test for mitrous acid with potentium indide starch paper, and diamonium selt with alcoholic, alkaline 2-naphthol.

Fodium carbonate sonohydrate was added to a 0.0477 sole solution of phenol-4-sulfonic acid (Page 6) until alkalinity to blue litsus was obtained. Iced (O to 9°C.), dispotized smiline, prepared above,

and sodium carbonate monohydrate, 7.5 gm., were added slowly and alternately to the rapidly stirred, chilled (0 to 9°C.), alkalina phenol-4-sulfonic acid over a 1-1/2 to 2 bour interval. The phenol-4-sulfonic acid was kept below 9°C. and alkalinity to litams was held during the addition. Diasonium salt addition was controlled, to avoid self coupling, by spot testing the phenol-4-sulfonic acid with alcoholic, alkalina 2-naphthol. A light colored test was maintained.

The mixture was held at 0 to 9°C. for 8 hours, with occasional stirring, filtered, and washed with 15 to 20 cc. of maturated sodium chloride solution. The red-brown cake, 15.5 gm., was dissolved in 400 cc. of boiling methanol and filtered. The methanol was replaced with acctone and the solution was concentrated to heavy crystals, chilled to 9°C., and filtered. 12.1 gm. of orange, sedium 2-phenyl-aso-phenel-4-sulfonate was obtained.

The cake filtrets was acidified to Congo Red with A.C.S. grade hydrochloric acid, saturated with sodius chloride, chilled to FC., and filtered. The filtrate was extracted twice with ethyl other and the cake was extracted with methonol. The combined other and methonol extracts, taken to dryness, yielded 1.6 ga. of clive green, 2-phenyl-aso-phenol-4-sulfonic hold.

Total yield calculated as 2-phenyl-aso-phenol-4-sulfonic soid was 12.8 gm., 96.7% of theoretical based on sulfanilic soid.

Identification tests on the product agreed with those in Beiletein (4).

# PREPARATION OF L.A'-DIAMINO-3-HYDROXY-DIFFESYL-6-SULFONIO ACID

The method of Whitmore and Revukas (28) was used, in the reduction of 2-phenyl-aso-phenol-4-sulfonic acid, as follows: Sodium 2-phenyl-aso-phenol-4-sulfonate, 5.0 gm. (0.0167 mole), see above, was dissolved in a mixture of 30 cc. distilled water and 80 cc. methanol in a Surgese-Parr hydrogenator (12). Raney nickel catalyst, Raney Catalyst Co., Chattanooga, Tenn., was activated by the method of Covert and Adkins (8) using methanol instead of ethanol. 9.0 gm. of the wet, activated catalyst in 20 cc. of methanol were added to the solution in the hydrogenator. 15 paig. hydrogen was introduced, through a 1.770 liter storage tank connected to a pressure gauge, and aritation was started.

In 20 minutes reduction was completed, as evidenced by no further drop in storage tank pressure. The tright red solution in the hydrogenator turned a pale yellow during reduction. The solution was transferred from the hydrogenator with methanol and the catalyst filtered off and washed well with methanol.

The filtrate turned a dark green, after stending 2 to 3 minutes. The cake was then extracted with 2% equeous, sodius hydroxide until a sample of the wash gave no precipitate when acidified, to litmus, with hydrochloric soid. Great care was taken to keep the catalyst wet with solvent, at all times, to prevent a fire, and it was dusped issediately into water after extraction.

The methanol and sodium hydroxide filtrates were combined and extracted with ethyl ether. These ether extracts, when dried over anhydrous calcium sulfate and concentrated to dryness, yielded a trace of brown tar. This tar showed no evidence of aniline and was not investigated further.

The extracted filtrete yielded 1.2 gm. of pink 4,4'-diamino-3-bydroxy-diphenyl-6-sulfonic ecid, after acidification to litaus with A.C.S. arade hydrochloric acid and chilling (0 to 5°C.) for 8 hours. More product, 2.0 gms., was isolated by concentrating the mother liquor, chilling (0 to 5°C.), and adding ethyl ether.

Total yield of 4,4'-dismino-3-hydroxy-diphenyl-6-sulfonic acid was 3.2 gas. 68.7% of theoretical. Identity was confirmed by microscopic comparison with 4,4'-dismino-3-hydroxy-diphenyl-6-sulfonic acid prepared from acidium 2-phenyl-aso-phenol-4-sulfonate using stannous chloride and hydrochloric acid (3).

A repeat reduction using a larger amount of renew nickel catelyst, 20.0 ga., on 5.0 gm. (0.0167 mole) of sodium 2-phenyl-aso-phenol-4-sulfonate, dissolved in 200 cc. of methanol only, gave the same results bensiding rearrangement and the production of 4,4' dismino-3-hydroxy-diphenyl-6-sulfonic acid, 2.7 gm., 57.8% of theoretical. The same reduction filtrate color change was noted. The ether extraction procedure was not used. Additional evidence for bensiding rearrangement was hydrogen consumption, measured by storage tank pressure drop, 1.2 moles per sole of aso compound.

## PREPARATION OF A-MITHO PHEND -(1 AZO 6)-PHENOL-4-SULFUMIC ACID

20N 
$$NH_2$$
  $HONO$   $20N$   $N:NC1$ 

BY HO SON  $N:NC1 + H_3OS$   $OH$   $2ON$   $N=N$ 

C.P. p-nitro aniline, 5.6 gm. (0.0406 mole), wer dissotized by the method of Sunder (22), using hydrochloric acid.

of phenol-4-sulfonic soid (Page 6) until alkalinity to blue liteus was obtained. The iced (O to FC.), dissotised p-nitro aniline and sodium cerbonate sonohydrate, 21.0 gm., were added slowly and alternately to the rapidly stirred, chilled (O to FC.), alkaline phenol-4-sulfonic acid over a 1-1/2 to 2 hour period. The phenol-4-sulfonic soid was kept below FC. and alkaline to blue liteus during the addition. Self coupling of the diamonium salt was kept to a minimum by controlling addition so that a light colored test was estimated when the phenol-4-sulfonic soid was spot tested on alcoholic, alkaline 2-naphthol.

The mixture was held below FC. for a hours, with occasional stirring, carefully acidified with A.C.S. grade hydrochloric acid to Congo Red, chilled to FC., and filtered. The dried orange or be yielded 15.4 gm. of ispure 4-nitro phenyl-(1 asc 6)-phenol-4-sulfonic acid. It was dissolved in boiling methanol and filtered. The methanol was replaced with ether and the solution was concentrated to heavy crystals, chilled to FC., filtered, and dried. 12.2 gm. of yellow-

orange, 4-mitro phenyl-(1 azo 5)-phonol-4-sulfonic acid was obtained, 90.16 of the theoretical based on p-nitro aniline.

Rentity: Calculated nitrogen, 13.0%. Found nitrogen, 12.9%. No reaction when titrated with sodium nitrite.

### PREDICTION OF A TITTO PRESENT. (1 AGO 6) - PRESENTA-CULTONIC ACID

$$2^{O_N} \longrightarrow 10^{O_N} = 10^{O_N} \longrightarrow 10^{O_N} \longrightarrow$$

contino phenyl-(1 aso 6)-phenol-4-sulfonic acid, 5.0 gm. (3.0135 mole), was dissolved in a solution of 1.85 gm. (0.0462 mole) ...3.3. grade sodium hydroxide pellets in 50 cc. distilled water, heated to 40°C. C.P. sodium hyposulfite, 6.3 gm. (0.0562 mole), was added, with stirring, and the solution color changed from a deep red to a yellow. He crystals appeared after chilling the alkaline colution — evidence that little or no p-nitro aniline or para phenylonedicaline was formed in the reduction. After acidification to Congo Red, with A.C.S. grade hydrochloric acid, the solution was carefully concentrated to crystals, chilled, and filtered. The dry, pink cake, 3.5 gms., was identified a cither ? -nitro-5-hydroxy-benzidine-6-sulfolds acid or 2.5-diamine-2 -hydroxy-diphenylamine-8 -sulfonic acid by nitrogen assay and sodium nitrite titration for anine groups. The calculated nitrogen agreed with the assay nitrogen for both compounds and no attempt was made to further establish identity.

Compound	Cole. E	Norma I.	Gan.	Yiold :: Theory
benzidine	12.93	15.00	0.94	10.7
dirhonylomine	14.2	14.3	0.85	18.8

Instability of nitro groups, in reducing notia, suggests that 2,5-diamino-2'-hydroxy-diphonylamino-5'-sulfonic acid is the more probable compound.

Because of this evidence that blocking the para position with a nitro group does not hinder rearrangement, the filtrate was not investigated further.

## PRUPARACICE OF NAPHTHOL-2-(1 AZO 6)-PHENOL-4-SULFOLIC ACID

A method similar to that used for Orange II (5) was used for the proparation of naphthol-2-(1 aso 5)-phenol-4-sulfonic acid.

2-Amino phenol-4-sulfonic acid, 98.9% pure, 9.9 gm. (0.052 note), was dissolved in a solution of 2.7 gm. (0.025 note) anhydrous sodium carbonate in 50 cc. distilled water and chilled to 15°C.

U.S.P. sodium nitrite, 3.7 cm. (0.055 mole), was discolved in 10 cc. distilled water and combined with the sulfonic acid solution. The whole was poured slowly into a rapidly stirred mixture of 10.6 cc. (0.125 mole) A.C.P. grade hydrochloric acid and 60 cm. ice. The mixture cave a positive test for excess nitrite with potassium iedide

starch paper. Piezonium salt formation was confirmed by spot test with alcoholic, alkaline 2-naphthol. A red-brown color was obtained.

c.7., alpha free, 2-maphthol, 7.2 (m. (0.000 mole), was dissolved in a warm solution of 7.5 (m. (0.183 mole) A.C.5. grade sodium hydroxide pellets in 60 cc. distilled water. The whole was then iced to 0°C., and 40 gm. ice was added. The rapidly stirred 2-maphthol solution was kept at 0 to 10°C. while the sulfonic acid diagonium solt was added over a 1-1/2 to 2 hour period. The 2-maphthol solution was kept alkaline to litual during the addition. After 2 hours, coupling was complete and the red-brown mixture was carefully acidified to Congo Red with A.C.5. grade hydrochloric acid. The thick slurry was heated on a hot plate, with stirring, to 40 to 60°C. At this point the foun disappeared and the solids settled to the bettom of the beaker. After chilling to 0 to 5°C., the mixture was filtered and the red-brown cake was washed once with a small portion of chilled, saturated sodium chloride solution and dried.

The yield of rod-brown, naphthol-2-(1 aso 6)-phonol-1-sulfonic acid was 16.9 gm., 98.20 of theoretical based on 2-naphthol. This raterial was reduced without further purification. Identity was established by the reduction products: 1-amino-2-naphthol and 2-amino phonol-4-sulfonic acid.

PROPARATION OF 1-ACTIO-2-LAPRING INDICOMINE AND 2-ACTIO PRESCRI-

$$\begin{array}{c}
\text{Na}_{2} \text{SO}_{2} \text{H} \\
\text{Na}_{2} \text{Na}_{2} \text{OH} \\
\text{SO}_{3} \text{H}
\end{array}$$

$$\begin{array}{c}
\text{OH} \\
\text{NH}_{2}
\end{array}$$

$$+ \qquad \begin{array}{c}
\text{OH} \\
\text{NH}_{2}
\end{array}$$

Maphthol-2-(1 aso 6)-phenol-4-sulfonic soid, 5.0 gs. (0.0145 sole). was dissolved in a AO to 50°C. solution of 3.2 gm. (0.080 mole) A.C.S. grade acdium hydroxide pellete in 28.8 cc. distilled water. C.P. sodium hyposulfite, 6.0 gs. (0.0285 mole), was edded countiously and the solution was quickly heated to 70°C., with stirring. Heating was continued until frothing subsided. The red solution turned white and meterial precipitated. After cooling to 20 to 25°C, and filtering, the cake was dissolved in a warm solution of 0.05 gm. C.P. stannous chloride in 23 cc. distilled water containing 1.8 cc. A.C.S. grade hydrochloric acid. The whole was stirred and heated until solution was complete. 2.9 cc. A.C.S. grade hydrochloric acid was added and the solution was chilled to O'C. and filtered. The white cake was air dried, below 35°C., on filter paper. 2.0 gm., 70.5% theoretical yield, of 1-amino-2naphthol hydrochloride was obtained. Identity was confirmed by exidising a sample to 1-2 naphthoguinens (decomposition melting point r 116°C.) with ferric chloride (6).

The filtrate efter acidification to Congo Red with A.C.S. grade hydrochloric acid, chilled (O to 5°C.) for 46 hours and filtered, yielded 1.9 ga., 69.3% theoretical yield of pale pink 2-amino phenol-4-sulfonic acid. Identity tests agreed with those in Deilstein (2) and microscopic comparison with 2-amino phenol-4-sulfonic acid substantiated composition.

# PRETIRETION OF NAPHTHOL-1-(L AZO 6)-PHENOL-L-SULPONIC ACID

$$_{\mathrm{H_{3}OS}}$$
  $_{\mathrm{OH}}$   $_{\mathrm{OH}}$   $_{\mathrm{OH}}$   $_{\mathrm{OH}}$   $_{\mathrm{OH}}$   $_{\mathrm{OH}}$   $_{\mathrm{OH}}$   $_{\mathrm{OH}}$ 

Naphthol-1-(4 and 6)-phenol-4-sulfonic and was prepared by coupling 9.4 cm. (0.050 mole) diasotized 2-amino phonol-4-sulfonic acid. (Page 15), with 6.9 gm. (0.048 mole) 6.5. 1-naphthol. The same procedure (Page 15) as for 2-naphthol was used for coupling and isolation of the product.

The 1-naphthol solution, at the start of diazonium palt addition, became a bright orange. This color turned a deep red toward the end of the addition. Additication produced a heavy, creamy precipitate which congulated when heated.

Yield of Green-brown, naphthol-1-(4 aso 6)-phenol-4-sulfonic acid was 9.1 gm., 52.8% of theoretical, based on 1-naphthol. This product was used directly in a reduction without further purification.

PROCEIGE OF HAPHTHOL-1-(4 AZO 6)-PHENOL-4-SULFUEL ACHO

5.0 gm. (0.0145 mole) of Eaphthol-1-(4 aso 5)-phenol-1-sulfonic acid was dissolved in a 40 to 80°C. solution of 5.2 gm. (0.080 mole)

A.C.S. grade sodium hydroxide pellets in 28.8 cc. distilled water.

C.S. sodium hyposulfite, 6.0 gm. (0.0285 mole), has added cautiously and the dark red solution was quickly heated to 70°C. and Jairred until frothing subsided. The color changed to an arange. The solution

diluent. After cooling to 20 to 25°C., the solution was filtered. During filtration the cake turned from a tan to pink. It was quickly transferred to a were solution of 0.05 ga. C.F. stannous chloride in 23 co. distilled water containing 1.8 cc. A.C.S. grade hydrochloric acid and heated to dissolve. The solution color was similar to that for 1-smino-4-naphthol hydrochloride (7), but the naterial did not completely dissolve. 5.6 cc. more A.C.S. grade hydrochloric acid was asked, the solution was chilled to O°C. and filtered. Again the cake color changed during filtration, from a tan to a pink. This color change may be observed when 1-smino-4-naphthol is prepared from Orange I (7). The yield, after drying below 35°C. on filter paper, was 0.5 ga. or 17.65 theoretical, presuming the material to be 1-amino-4-naphthol hydrochloride. Positive identification was not made because of the low yield.

Acidification of the filtrate to Congo Red with hydrochloric acid and chilling (O to 9°C) overnight yielded no material.

After consideration of the low coupling yield and the extremely emall reduction yield, investigation of the 4-emo-1-nephthol was terminated.

### DLAZOZIZATION OF 1-AUNIO-2-NAPHTROL

When applied to 1-amino-2-maphthol, the method using sodium scatte (15) proved to be most convenient and easily managed.

An attempt to acetylate, using the method of Krosber and Jagerspacher (16) was unsuccessful.

Dissotisation in aqueous copper sulfate, recommended by Sandmayer (20) and Modgson and Sirtwell (14), yielded a gum which was useful only when dissolved in acetic acid.

The sodium acctate procedure was used as follows: Analytical grade 1-amino-2-nephthol hydrochloride (5), 2.0 gm. (0.0102 mole), was dissolved, by warming in a solution of 1.02 gm. (0.0124 mole) anhydrous sodium acctate in 10.2 cc. (0.1772 mole) acctic acid. After cooling to 20 to 25°C., 0.85 gm. (0.0123 mole) U.S.P. sodium nitrite was slowly added. A dark red-violet solution of disso-oxide formed. This solution coupled slowly with elecholic, alkaline 2-nephthol, on a spot test, to give a blue-black color.

## COUPLING OF 1-AMENO-2-MAPHTHE. DIAZO-OXIDE WITH PHENOL-A-SULFONIC ACID

All attempts to couple 1-amino-2-maphthol diago-oxide with phenol-4-sulfonic acid failed. Solutions of phenol-4-sulfonic acid in dilute acetic acid and aqueous, alcoholic, potessium hydroxide (alkaline to brilliant yellow) were tried. Solutions were allowed to stand for as long as 3 days. There was no visible color change in the solution and positive coupling tests to 2-maphthol were obtained.

Acidification of all solutions, at the end of 3 days, with A.C.S. grade hydrochloric acid produced a small amount (0.5 to 1.0 gm.) of red-violet material. This material was not soluble in sodium hydroxide

and sould not be reduced with sodium hyposulfite. This proved that it was not the desired production, nambthol-2-(1 aso 6)-phenol-4-sulfonic soid (Page 13 and 14) which ensily reduced. No further identification was made.

De Jonge and Dijkstra (10) investigated irradiation decomposition of o-hydroxy dissentus compounds, in said solution. Sed dyes were isolated which were proved to be the dissentus salt coupled to its decomposition product, indens and indens disarboxylic acid. The red dye isolated here night be that mentioned by De Jonge and Dijkstra.

DIAZOTIZATION AND COUPLISO OF 1-ASUNO-2-MAPRIMOL-4-SULPONIC ACID

$$\begin{array}{c}
\text{NH2} \\
\text{OH} \\
\text{SO_3H}
\end{array}$$
+ CuSO<sub>4</sub>
- NaNO<sub>2</sub>
- OH
- SO<sub>3</sub>H

l-amino-2-naphthol-4-sulfonic acid (9), 2.45 gm. (0.0102 mole), was dissotized in the presence of copper sulfate (25). The yellow-grade dissonium compound was added to a solution of 0.0133 mole phonol-4-sulfonic acid (Page 6) containing enough sodium carbonate to make it just alkaline to brilliant yellow.

The solution, after standing at 10°C. for 3 days, was filtered; and 2.45 gs. material was isolated and identified as the original diamo-oxide, by appearance and color when coupled to alkaline 2-naphthol.

Another reaction tried with an alkaline, methenol solution of phenol-4-sulfonic acid gave the same result, no coupling.

COUPLING OF 2-ANIMO PRESSA-4-CULTURING ACID -- 2-mains phenol-4-sulfonic acid, 98.9% pure, 9.9 gm. (0.052 mole), was dissocized as previously described (Page 15).

A 0.052 mole solution of phenol-4-sulfonic acid (Page 6) was alkalized with 6.0 gm. A.C.S. grade sodium hydroxide pellots, and chilled to 0 to 9°C. The solution of dissotized 2-amino phenol-4-sulfonic acid was added. During addition, alkalinity to liteus was maintained and the temperature of the combined solution was kept below 9°C. Very little color change was noted after combination.

At this point the solution was split into two parts and methanol equal to twice the volume was added to one half. Both parts were then allowed to stand at 0 to 9°C. for 4 days.

A coupling test on the undiluted half, with alkaline 2-maphthol, indicated that the original amount of diamonium salt was still present and the solution was stored as heldre for 3 days acre. After this second period, a test showed that diamonium salt was still present and coupling had not taken place. This method for coupling was not investigated further.

After 4 days, the methenol diluted half showed no signs of diamonius salt, even though the color was the same as before stending. No crystals appeared when the solution was acidified to Congo Red, with A.C.S. grade hydrochloric acid, heated until all methanol was removed, and chilled to 8 to 9°C.

A reduction on the amber solution without isolating the azo dye, if formed, was tried. A.C.S grade sodium hydroxide pellets were added

until elkelinity to liteue was obtained. Then 6.0 gm. wore

(0.150 mole) elkeli was added; the solution was heated to 50 to 60°C.

12.0 gm. (0.057 mole) C.P. sodium hyposulfite was added. The solution was stirred until no more color change was noted, filtered, scidified to Congo Red with A.C.S. grade hydrochloric soid, chilled to 0 to 9°C. overnight, and filtered.

Material, 5.4 gm., isolated was identified as a compound containing no amine groups, by titration with nitrous acid. Amines were not found in the acid filtrate.

Coupling of 2-amino phenol-4-sulfonic acid diagonium salt with phenol-4-sulfonic acid was therefore considered negligible.

PREPARATION OF MAPRIMIL-(2 AZO 6)-PHENOL-4-SULFONIC ACID

2-maphthylamine, 5.85 gm. (C.OLOS mole), was dissotized a coording to Scheides (26) procedure. The solution gave a brilliant red coloration when spot tested with elkaline 2-maphthol.

This dissentum selt was added to a chilled (0 to 9°C.), stirred, solution of 0.0408 sole phenol-4-mulfonic soid (Page 6) containing 50.7 gm. (0.0408 sole) sodium carbonate monohydrate. After standing 48 hours at 0 to 9°C., coupling was complete as determined by spot

testing with alkaline 2-naphthol. The solution was acidified to Congo Red with A.C.S. grade hydrochloric acid, chilled to O to 5%C., and filtered.

Green-brown Maphthyl-(2 aso 5)-phenol-4-sulfonic acid, 12.8 gs., 95.5% of theoretical, was obtained. The almost colorless filtrate was discarded.

Identity was established, see below, by reduction to 2-nephthylamine and 2-smino phonol-4-sulfonto soid. Furity was estimated at 79%.

Crystallisation from methanol can be affected by dissolving, filtering, concentrating to heavy crystals, diluting with ethyl other, and chilling. A 78% yield of purified, yellow-green Haphthyl-(2 and 6)-phenol-4-sulfonic acid was obtained by such a procedure.

For reduction purposes, a better yield was obtained by working directly with the crude dys.

# RESERVETION OF HAPPERIL-(2 AZO 6)-PHENOL-A-SULFUFIC ACID

Crude Maphthyl-(2 aso 6)-phenol-4-sulfonic acid, prepared above, 10.0 gm. (0.0305 mole), was dissolved in a mixture of 5.3 gm. (0.13%2 mole) A.C.S. grade sodium hydroxide pallets in 120 cc. distilled water by heating to 35 to 50°C. The red solution was filtered and 20 to 30 cc. distilled water was used to wash the cake and transfer the filtrate to

a clean beaker. The dried cake weighed 2.1 gm. (melting point \* 184 - 185°C.) indicating that the crude dye was approximately 79% pure and the charge was actually 7.9 gm. (0.024 mole).

C.P. sodium hyposulfite, 14.1 gm. (0.067 mole), was added, with stirring, to the warm filtrate, 35 to 45°C. The solution thickened and turned gray. Shen forming subsided, the solution was chilled, 0 to 5°C., and filtered. The insoluble, 2-nephthylamine, oxidized when allowed to stand in the air and was converted to the more stable hydrochloride.

The light pink cake was quickly dissolved in a hot, 40 to 60°C., mixture of 5.2 cc. A.C.S. grade hydrochloric soid in 80 cc. distilled water. An equal volume of aqueous, seturated sodium sulfate solution was added and the whole was chilled, 0 to 5°C., and filtered.

2.6 gm. of 2-naphthylamine hydrochloride, 64.8% of theoretical based on the corrected 0.024 mole charge of asc dye, was obtained. Identity was established by scetylation to form N-acetyl-2-naphthylamine, (melting point melting point mes run with known N-acetyl-2-naphthylamine. Conversion of the 2-naphthylamine hydrochloride to 2-naphthylamine by mining in water and alkalizing to phenolphthelein, yielded material that had a melting point of 110-111°C. No melting point depression was exhibited when a mixed melting point was run with known 2-naphthylamine.

The reduction filtrate, when acidified to Congo Hed with A.C.S. grade hydrochloric acid, concentrated to 75 cc., chilled (C to 9C.)

for 48 hours, and filtered, yielded 2.1 gm. of pale pink, 2-smino phenol-4-sulfonic acid, 46.1% of theoretical. Identity tests agreed with those in Beilstein (2) and microscopic comperison with 2-smino phenol-4-sulfonic sold substantiated composition.

Reduction of 10.0 gm. (0.0305 male) Nephthyl-(2 exc 6)-phenol-4-sulfonic soid, purified by methanol crystellization, (Page 22), gave 3.0 gm., 68.8% theoretical, 2-maphthylamine and 3.0 gm., 52.2% theoretical, 2-main phenol-4-sulfonic soid.

# PREPARATION OF MAPRICIL-(1 ero 6)-PHENOL-4-GULFONIC ACID

$$\begin{array}{c|c}
 & \text{HONO} \\
 & \text{HC1} \\
 & \text{N:NC1} \\
 & \text{HO} \\
 & \text{SO}_{3H}
\end{array}$$

$$\begin{array}{c|c}
 & \text{N:NC1} \\
 & \text{N:NC1} \\
 & \text{SO}_{3H}
\end{array}$$

1-naphthylamine, 11.7 gm., (0.0653 mole), was dissotised according to the procedure of Basberger (1). Dissotisation temperature was held at 15 to 18°C. The reaction was slow at lower temperatures. The solution give a red-purple color when spot-tested with alkaline 2-naphthol.

This diamonium solution was added to a rapidly stirred, chilled (O to 9°C.) solution of 0.0952 sole phenol-4-sulfonic acid, (Page 6), containing 115 gm. (0.926 sole) sodium carbonate monohydrate. After standing 48 hours at O to 9°C., coupling was complete as indicated by

a negative spot test with alkaline 2-neghthal.

The solution, after ecidification to Gongo Red with A.C.S. grade hydrochloric acid, was chilled (O to 5°C.) and filtered. Red-brown, crude Naphthyl-(1 aso 6)-phenol-4-sulfonic acid, 19.8 gm., 73.9% of theoretical based on 1-naphthylamine, was obtained. The almost colorless filtrate was discarded.

Identity was established by reduction to 1-maphthylamine end 2-amino phenol-4-sulfonic acid, see below. Purity was estimated at 58%.

## HEDNOTION OF MAPPIPHYL-(1 AZO 6)-PHENOL-L-SOLPONIC ACID

Crude Maphthyl-(1 aso 6)-phenol-4-sulfonic acid, prepared above, 10.0 gm. (0.0305 mole), was dissolved in 120 cc. distilled water containing 5.3 gm. (0.132 mole) A.C.S. grade sodium hydroxide pallets by heating to 40 to 50°C. The insoluble, 4.2 gm. (malting point : 207 - 208°C.) indicated that the crude aso dye was approximately 58% pure and the reduction charge was corrected to 5.8 gm. (0.0177 mole).

C.F. sodium hyposulfite, 14.1 ga. (0.067 mole), was added, with stirring, to the filtrate at room temperature. The red solution turned pink. It was chilled (0 to 9°C.), filtered, and the cake was dissolved in a mixture of 2.6 cc. A.C.S. grade hydrochloric acid in 40 cc. of distilled water.

1-naphthylemine oxidized quickly in air and was recovered as the more stable hydrochloride as follows:

An equal volume of aqueous, saturated sodium sulfate solution was added, the mixture was chilled to 0 to 500, and filtered. 2.5 gm. of 1-naphthylamine hydrochloride was obtained, 71.50 of theoretical.

Identity was established by converting a sample to 1-naghthylactine by sludging in vator and alkalizing to ghenolphthalein. The
insoluble 1-naphthylemine melted at 13 - 50°C, and no change in
melting point was observed when a sample was mixed with known
1-naphthylamine.

The reduction filtrate, when acidified to Congo Red with A.J... grade hydrochloric acid, concentrated to half volume, chilled (0 to 5°C.) for 48 hours and filtered, yielded 1.2 gms. of pale pink: 2-amino phenol-4-sulfonic acid, 55.85 of theoretical. Identity tests agreed with those in Beilstein (2) and microscopic comparison with 2-amino phenol-4-sulfonic acid substantiated composition.

## EXPERIMENTAL RESUMS

A total of six coupling reactions were run. Each involved too separate processes: (1) preparation of the diazonium salt, and (2) coupling of the diazonium salt. The various compounds propared, methods, and yields are summarized in Table I. Page 27.

In addition to the dissotisations listed in Table I, Page 27.

others were carried out and were abandoned because they would not couple with phenol-4-sulfonic acid. Their preparation: . properties are presented in Table II. Page 28.

Reduction of the ase compounds listed in Table I. Page 27.

27

Table I -- Aso Compounds Prepared by Coupling Reactions

	Dissetized Coaround	Diazoties <u>Procedu</u>		Coupled <u>To</u>		Product	Yiold
ć	Ani <b>line</b>	direct	(23)	Phenol-4-sulfonie	bles	2-phenyl-aso-phenol- 4-sulfonio soid	96.7%
	p-nitro eniline	direct	(23)	Phenol-4-sulfonic	acid	4-nitro phenyl-(1 eso 6)- phenol-4-sulfonic acid	93.1
	2-amino phenol- 4-sulfonic soid	inverted	(22)	2-saphthol		maphilial-2-(1 aso 6)- phonol-4-sulfonic acid	98,2
	2-saino phenol- 4-sulfonio ecid	inverted	(22)	1-naphthol		phonol-4-sulfenic sold	52.8
	2-muphthylesine	direct	(26)	phenol-4-sulfenie	acid	maphthyl-(2 aso 6)- phenol-4-sulfonic soid	75.5
	1-naphthylamina	direct	(1)	phonol-4-sulfonio	acid	mephthyl-(1 aso 6)- phenol-4-sulfonic acid	42.8

X Y:

TABLE II - Diazonium Compounds that do not Couple with Phenol-4-sulfonic Acid

Diamotimed Compound	Diesotisetion Procedure	Properties
l-amino-2- naphthol	sodina soctate inhibited (15)	red-violet solution in acetic acid. Violet gum in water. Couples to alkaline 2-naphthol to produce blue-black dye.
l-amino-2- sephthol	acetylation of hydroxyl (16)	did not diamotime.
l-amino-2- naphthol-4- sulfonic seid	copper sulfate inhibited (25)	yellow-gray folid. HCl insol. Easily couples with alkaline 2-naphthol to produce purple dyo.
2-amino phemoi- 4-sulfonis acid	invorted (22)	white solid. MSI insol. Easily sourles to alkaline 2-maphthol to produce a red-brown dye, Page 14.

Table III -- Reduction of Aso Derivatives of Phenol-4-Sulfonic Soid

áso Compound	Reduction <u>Setion</u>	Reaction	Product	_Lield_
2-phenyl-mac-phenol- 4-sulfonie acid	Raney nickel catalyst	Pensidiae rearrangement	4, 4' diamino-3- hydroxy-diphenyl- 6-culfonte acid	68.7%
4-nitmo phenyl-(1 azo 6) phenol-4-sulfonie scid	- Alkaline sodium hyposulfite	semidine rearrangement	probably 2,5- diamino-2'-bydroxy- diphenylamino-6- sulfonic ecid	18.8
Naphthol-2-(lese 6)-	Alkaline	fiseion	1-smino-2-maybthol	70.5
sulfonic scid	hyposulfite		2-amino phenol-4- sulfonis soid	69.3
Waphthol-1-(4 aso 6)-	Alkalice sodium	fission	1-amino-4-maphthol	17.6
sulfonic acid	hypoculfite		nothing clse isolated	
Mephthyl-(2 aso 6)- phonol-4-sulfonic	Alkaline sodice	fission	2-nephthylanine	64.8
aci6	hyposulfits		2-emino phonol-4- mulfonio edid	46.1
Wephthyl-(1 aso 6)	Alkaline	fission	1-aephthylosine	72.5
serg	phoenstire		2-amino phenol-4- sulfonic soid	35.8

others formed semidines, and others split and gave the desired smines. Heduction results are summarized in Table III, Fage 29.

#### DISCUSSIVA

There were two problems to be overcome before a satisfactory solution to this investigation could be resched: (1) selection of an amine that could be dissotised and coupled to phenol-4-sulfonic acid, and (2) thereby produce an azo compound that would split, when reduced, and not rearrange.

The problem was further complicated when it because evident that phenol-4-sulfonic acid was a weak acceptor and required a strong diamonium salt compler.

DIAZOTIZATION -- Four general techniques were used for dissotisation in this investigations direct method, inverted method,
sodium acetate inhibited method, and copper sulfate inhibited
method. Each procedure reference is listed, Table I, Page 27
and Table II, Page 28; and directions as given in the literature were
used, except for slight medifications as were deemed necessary.

The direct method was applied to those smines, smiline, pmitro smiline, and the mephthylamines, that were strongly basic and whose salts were soluble in aqueous acid.

The sulfonic soids, except for 1-amino-2-naphthol-4-sulfonic acid, were diszotized by the inverted method which is designed for those amines which are weakly besic or whose salts are almost insoluble in aqueous soid.

The 1-emino-2-nephthole were dissotized under inhibited

conditions to prevent the nitrous acid from oxidizing them to maphthoquinous (13). Fodius acetete and copper sulfate were used successfully as inhibitors, but a "pre-acetylation of the hydroxyl group" recommended by Kroeber and Jagerspacher (16) failed.

<u>COULDE</u> -- The coupling shility of the various diszonius compounds, in squeous solution sikaline to red litems, with phenol-4-sulfonic acid is:

aniline > p-nitro amiline > 2-naphthylamine > 1-naphthylamine

This order is based on yields from the coupling reaction,
Table I, Page 27, presuming that side reactions, such as self
coupling and decomposition are negligible. Coupling yields varied
from 98% for the aniline diagonium compound to 43% for the 1-naphthylamine diagonium compound.

Phenol-4-sulfonic soid is a week acceptor, compared to 2-maphthol, and coupling may not take place or proceed to a receonable yield unless a strong coupling diasonium selt is used.

The diasonium compounds listed below would not couple with phenol-4-sulfonic soid in acid or alkaline solution, see Table II., Page 28, and are rated on ability to couple with 2-nephthol:

2-amino phenol-4-sulfenic ecid > 1-amino-2-naphthol-4-sulfenic ecid >>> 1-amino-2-naphthol

1-amino-2-naphthol would have been an excellent solution to this investigation if it suld have been coupled to phenol-4-sulfonic acid, but it is such a poor coupler that its reaction with 2-naphthol,

s very strong acceptor is slow. The aso dye, prepared in reverse by coupling dissotised 2-amino phenol-4-sulfonic seid with 2-naphthol, reduced easily and split into 1-amino-2-naphthol and 2-amino phenol-4-sulfonic acid, see Table III, Page 29.

<u>REMICTION</u> -- Products and techniques of reduction are susserized in Table III, Page 29.

Whiteere and Revukes (28) ennounced fission of ase benzens, a compound which may be reduced to benzidine, by reduction with rancy nickel catalyst and bydrogen under pressure. Their procedure was applied in this investigation to 2-phenyl-ase-phenol-4-sulfonic acid without success. The same bensidine rearrangement produced in reductions with stannous chloride and bydrochloric acid was obtained, with 69% yield.

Except for 2-phenyl-ase-phenol-4-sulfonic edid, reductions sere performed by adding sodium hypomulfite to the strong mikaline solution of the ase dye. Weakly basic or neutral solutions may produce bydrama compounds (18).

Blocking the 4-position with a nitro group did not promote fission and 4-nitro phenyl-(1 see 6)-phenol-4-sulfenic acid did not split when reduced. Positive identification was not completed, but either a beneiding or semiding was produced. A semiding is more likely since Jacobson (17) reported that semidines resulted shen 4 and 4'-hydrazo bensens positions were occupied with brose, ethoxy, iede, or methyl groups.

Fission of 1-aso derivatives of 2-naphthol (5) and 2-naphthol scotate (11) has been reported. This investigation of naphthol-2-

(1 ezo 6)-phenol-4-sulfonic acid describes a similar fission with a 70% yield. As was previously indicated, Fage 31, this dye would be an excellent source material for 2-amino phenol-4-sulfonic acid if a method for coupling the 1-amino-2-naphthol diszonius compound to phenol-4-sulfonic acid could be developed.

A new synthesis for 2-smino phenol-4-sulfonic soid is described:
Coupling of the dissonium salt of either 1- or 2-naphthylamine to
phenol-4-sulfonic soid, followed by reduction of the resultant azo
compound. Of the two smines, 2-naphthylamine is preferable. It is
higher melting and more stable, and gave a 76% higher coupling yield.
The 1- and 2-naphthylamine derivatives were split in alkaline sodium
hypomulfite to produce 2-smino phenol-4-sulfonic acid, 46% and 36%
yields respectively, and the corresponding 1- and 2-naphthylamine,
65% and 72% yield respectively.

### SUMME

Procedures for the preparation and reduction of some 2-azo derivatives of phenol-4-sulfonia acid are given.

Phenol-4-sulfonic seid is a weak acceptor in a coupling reaction and requires a strong dissonium selt coupler.

Order of coupling of the various diagonius selts studied, with phenol-4-sulfonic acid is proposed:

antline p-mitro entline 2-maphthylemine

1-naphthylasine

The disso-exides studied did not couple with phenol-4-sulfonic acid. Their ability to couple with 2-naphthol is proposed:

2-amino phenol-4-sulfonic soid > 1-amino-2-naphthol-4-sulfonic soid >> 1-amino-2-naphthol

Raney mickel catalyst and hydrogen under pressure reduction of 2-phenyl-aso-phenol-4-sulfonic acid produced a bensidine, 4-4'-diamino 3-hydroxy-diphenyl-6-sulfonic acid.

Slocking the 4-position with a nitro group did not promote fission and 4-nitro phenyl-(l see 6)-phenol-4-sulfonic soid did not split when reduced in alkaline sodium hyposulfite. A semidine or bensidine was produced.

Reduction of Naphthyl-(2 ase 6)-phenol-4-sulfonic acid and Naphthyl-(1 ase 6)-phenol-4-sulfonic acid in alkaline sodius by posulfite is presented as a heretofore undescribed sethod for the preparation of 2-asino phenol-4-sulfonic acid. These ase derivatives can be prepared by coupling the naphthylamine diagonium salt with phenol-4-sulfonic acid.

## LITERATURE CITED

- 1. Bamberger, Ber., 55, 3383-92 (1922).
- 2. Beiletein, Bandbuch der Organischen Chemie, XIV, System 1926, (614), 1931.
- 3. Ibid., (653), 1931.
- 4. Idde., XVI, System 2158, (293), 1931.
- 5. Elatt, A.H., "Organic Synthesis", vol. 2, p. 35, John wiley & Sons, London (1941).
- 6. Ibid., vol. 4, p. 430, John Wiley & Sons, London (1934).
- 7. Ibid, p. 39.
- 8. Covert, L.W., and Adkins, H., J. As. Chem. Soc., 54, 4116 (1932).
- 9. Cummings, Hooper, and Wheeler, "Systematic Organic Chemistry", p. 322, D. Van Hostrand Co., New York (1931).
- 10. De Jonge and Mjkstra, R.c. Trav. Chim., 67, 328-43 (1948).
- 11. Ghigi, E., Melv. Chim. Acta., 23, 426-30 (1940).
- Gillman, H. "Organic Synthesis"., vol. 1, 1st. ed., p. 53,
   John Wiley & Sons, New York (1932).
- 13. Grandwougin and Michel, Ber., 25, 972-85 (1892).
- 14. Hodgeon, H.H., and Hirtwell, S., J. Chem. Soc., 321-3 (1943).
- 15. Ibid., 539-40 (1944).
- 16. Erosbur and Jagurspacher, U.S. Patent 797, 441 (May 23, 1905).
- 17. Jacobson, P., Ann., 427, 142-221 (1923).
- 18. Peerl, 1.A., J. Org. Chem., 10, 205-210 (1945).
- 19. Robinson, H., J. Chem. Soc., 220 (1941).
- 20. Sandmeyer, U.S. Patent 793, 743 (Sept. 30, 1904).

- 21. Saunders, K.H. "The Arcantic Diaso-Compounds and their Technical Applications", 2nd. ed., Edward Arneld & Co., London (1949).
- 22. Ibid., p. 10.
- 23. Ibid., p. 6.
- 24. Ibid., p. 28-31.
- 25. Ibid., p. 30.
- 26. Schold, Sor., M., 1813 (1901).
- 27. Shrave, R.H., "Dyes Classified by Intermediates", P. 60-1, Chemical Catalogue Co., (1922).
- 28. Whitmore, W.F., and Revukes, A., J. Am. Chem. Soc., 52, 1500-3 (1937).